

ISOPROPYL ALCOHOL- isopropyl alcohol liquid

BuckAirways Inc.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Active Ingredient

Isopropyl Alcohol 91%.....First aid antiseptic

Uses

helps prevent risk of infection from:

- minor cuts
- burns
- lacerations
- burns

Warnings

For External use only. Internal ingestion will result in gastric problems.

Flammable

- keep away from fire, flame, spark, heat, electric

Ask a doctor before use

for deep lacerations, puncture wounds, animal bites, or any serious burns

When using this product

- avoid from contact with eyes
- do not apply over large areas of body
- do not use longer than 1 week unless instructed by a physician

Stop use and ask a doctor if

condition persists or gets worse

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center immediately.

Caution

Fumes created by Isopropyl Alcohol can be irritating to skin, eyes, or respiratory system. Do not apply if irritation is present on skin or continued irritation develops. Avoid from applying to eyes or mucous membranes. Do not inhale this product.

Directions

- clean the affected area
- apply a small amount of product on the area 1 to 3 times each day
- may be covered with a sterile bandage following application
- let dry prior to being bandaged

Other Information

- does not contain, nor is intended as a substitute for grain or ethyl alcohol

Inactive Ingredient

water

75396-071-18 LABEL PANEL

cln:™
IPA

91%

ISOPROPYL ALCOHOL

128 FL OZ
3.79L

Topical Cleansing Agent

Antiseptic & Sanitizer

Drug Facts

Active ingredient	Purpose
Isopropyl Alcohol 91%	First aid antiseptic
Uses: Helps prevent risk of infection from: • minor cuts • burns • lacerations • bites	
Warnings For External use only. Internal ingestion will result in gastric problems. Flammable: Keep away from fire, flame, spark, heat, electric. Ask a doctor before use for deep lacerations, puncture wounds, animal bites, or any serious burns.	
When using this product, avoid from contact with sensitive not apply over large areas of body with not use longer than 1 week unless instructed by a physician.	
Stop use and ask a doctor if condition persists or gets worse.	
Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center immediately.	
Caution: - Fumes created by Isopropyl Alcohol can be irritating to skin, eyes or respiratory system. Do not apply if irritation is present on skin or continued irritation develops. Avoid from applying to eyes or mucous membranes. Do not inhale this product.	
Directions: - Clean the affected area. Apply a small amount of product on the area 1 to 3 times each day until it is covered with a sterile bandage following application. Let dry prior to being bandaged.	
Other information: does not contain, nor is intended as a substitute for grain or ethyl alcohol. Irritates liquid water	

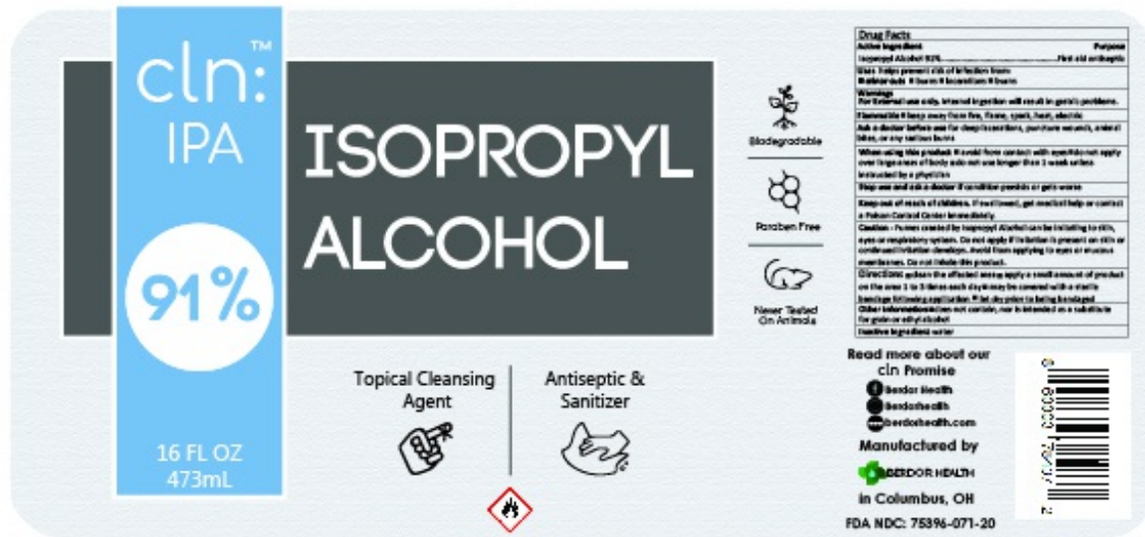
Read more about our cln Promise

BERDOR HEALTH
in Columbus, OH

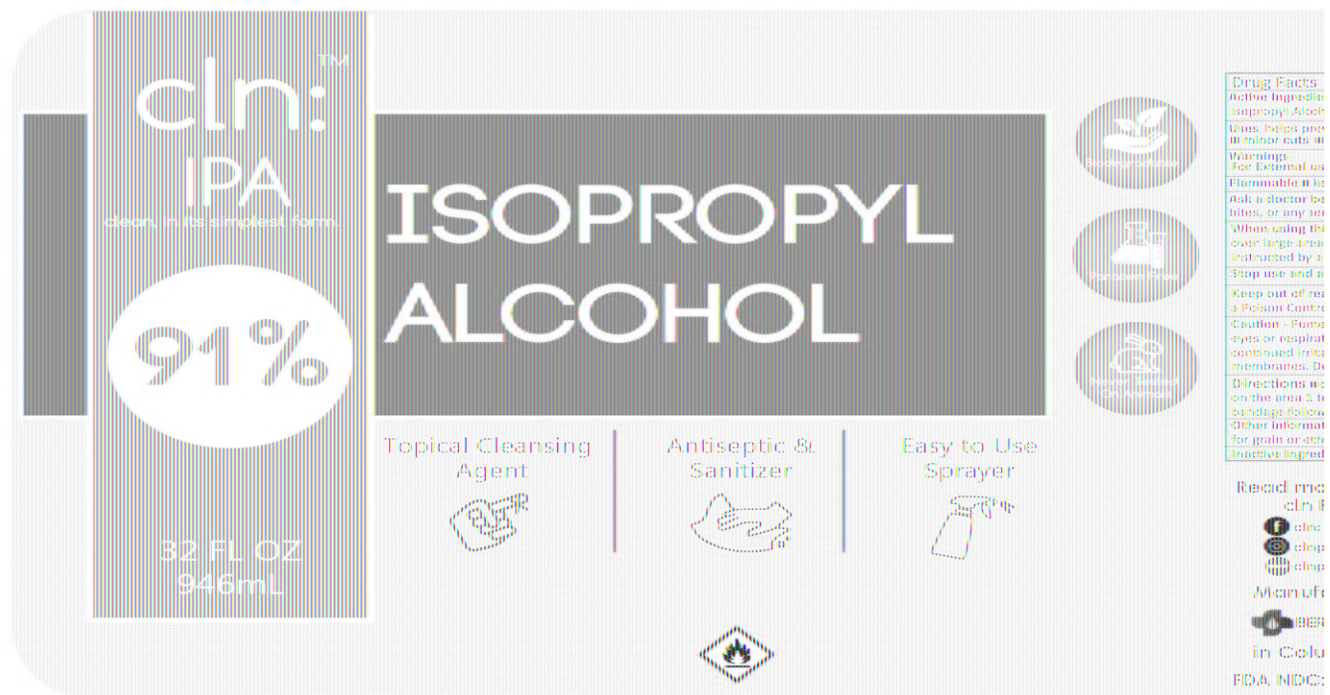
FDA NDC: 75396-071-18

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75396-071-20 LABEL PANEL



75396-071-21 LABEL PANEL



ISOPROPYL ALCOHOL

isopropyl alcohol liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:75396-071
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ISOPROPYL ALCOHOL (UNII: ND2M416302) (ISOPROPYL ALCOHOL - UNII:ND2M416302)	ISOPROPYL ALCOHOL	91 mL in 100 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:75396-071-18	3785 mL in 1 JUG; Type 0: Not a Combination Product	07/18/2020	
2	NDC:75396-071-20	473 mL in 1 BOTTLE; Type 0: Not a Combination Product	12/12/2020	
3	NDC:75396-071-21	946 mL in 1 BOTTLE; Type 0: Not a Combination Product	01/18/2021	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	07/17/2020	

Labeler - BuckAirways Inc. (117488255)**Registrant** - BuckAirways Inc. (117488255)**Establishment**

Name	Address	ID/FEI	Business Operations
BuckAirways Inc.		117488255	manufacture(75396-071)

Revised: 1/2021

BuckAirways Inc.